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(54)	FLOWARLE POLYAMIDES WITH HYPERBRANCHED	(30)	Foreign Applica	tion Priority Data

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- (57) ABSTRACT
- Thermoplastic molding compositions, comprising A) from 10 to 99% by weight of at least one thermoplastic polyamide,
- B) from 0.01 to 50% by weight of
- B1) at least one highly branched or hyperbranched polycarbonate with an OH number of from 1 to 600 mg KOH/g of polycarbonate (to DIN 53240, Part 2), or
- B2) at least one highly branched or hyperbranched polyester of A_xB_y type, where x is at least 1.1 and y is at least 2.1, or a mixture of these,
- C) from 0 to 60% by weight of other additives, where the total of the percentages by weight of components A) to C) is 100%.

FLOWABLE POLYAMIDES WITH HYPERBRANCHED POLYESTERS/POLYCARBONATES

[0001] The invention relates to thermoplastic molding compositions, comprising

[0002] A) from 10 to 99% by weight of at least one thermoplastic polyamide,

[0003] B) from 0.01 to 50% by weight of

[0004] B1) at least one highly branched or hyperbranched polycarbonate with an OH number of from 1 to 600 mg KOH/g of polycarbonate (to DIN 53240, Part 2), or

[0005] B2) at least one highly branched or hyperbranched polyester of $A_x B_y$ type, where x is at least 1.1 and y is at least 2.1,

[0006] or a mixture of these,

[0007] C) from 0 to 60% by weight of other additives,

where the total of the percentages by weight of components A) to C) is 100%.

[0008] The invention further relates to the use of the inventive molding compositions for production of fibers, of foils, or of moldings of any type, and also to the resultant moldings.

[0009] Pobyent/conties are usually obtained from the reaction of alcohols with phospings, or from transentification of alcohols or of phenols, using disklyl or disrylen/nontees. Arcunizie polyven/noutes are important insubstituly and are prepared, by way of example, from bisphenols, whereas alphalic polyven/noutes are less important in terms of alphalic polyven/noutes are less important in terms of Kanstsunf-Handhushi [Plasicise Handhood] Vol. 31, Polycurounts, Polyvenize, Polyvenize, Gallouseester [Polyvenboartes, polysecatis, polyvenize, cellulose estens]. Carl-Hanner-Verlag, Munich 1992, pages 118-119.

[0010] The aliphatic polycerbonates described are generally linear or else have a structure with a very small degree of branching. For example, U.S. Pat. No. 3,305,000 describes the use of solid linear polycerbonates whose molecular weight is above 15 000 dalton as plasticizers for polyvinyl polymers.

[0011] Low-molecular-weight additives are usually acked to thermoplastics in order to improve flowability. However, these additives have very limited effectiveness because, for example, the fall-off in mechanical properties becomes unacceptable when the amount added of the additive is increased.

[0012] High-functionality polycarbonates of defined structure have been known only for a short time.

[9013] S. P. Ramard and N. J. Davis, J. Am. Chem. Soc. 2000, 122, 11729 describe preparation of perfectly branched dendrimeric polycarbonates via reaction of carbonyfbiamidazole, as phosgene-analogous compound, with histydrorycitylamino-2-propanol. Synthese to give perfect dendrimers have four stages and are therefore expensive and not very suitable for industrial scale-up.

[0014] D. H. Bolton and K. L. Wooley, Macromolecules 1997, 30, 1890 describe the preparation of high-molecularweight, high-rigidity hyperbranched aromatic polycarbonstes via reaction of 1,1,1-tris(4'-hydroxy)phenylethane with carbonylbisimidazole.

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[0015] Hyperbranched polycarbonates may also be prepared as in WO 98/90453. In the process described there, triols are again reacted with carbonylbisimidazos. The first product is imidizabilides, and these are then further reacted intermolecularly to give the polycarbonates. The method mentioned gives the polycarbonastes in the form of colorless or pale yellow nubbery products.

[9016] The syntheses mentioned giving highly branched or hyperbranched polycarbonates have the following disadvantages:

[0017] a) the hyperbranched products either have high melting point or are rubbery, and

[9018] this markedly restricts subsequent processability.
[9019] b) imidazole liberated during the reaction has to be removed from the reaction mixture in a complicated

[0020] c) the reaction products always comprise terminal imidazolide groups. These groups are labile and have to be converted into, for example, hydroxy groups by way of a subsecuent step.

[9021] d) carbonyldiimidazole is a comparatively expensive chemical which greatly increases raw material costs.

[9622] WO-97/45474 discloses themoplastic compositions which comprise dendrimeir polyesters in the form of an Ally molecule in a polyester. Fire, a polyhydric alcohol produced in a polyester. Fire, a polyhydric alcohol and produced in produced in polyhydric alcohol Ally multicolate in given a dominantia polypoper. That courprises only OII functionalities at the set of the chair. Disadvantages of these mixtures are the high glass trunsition temperature of the dominimier to polyester, the compartively complicated preparation process, and especially the poor solubility of the dominimier is the polymanic matrix.

[9023] According to the teaching of DE-A101 32 928, the incorporation of branching secuts of this type by measure compounding and solid-phase post-condensation improves mechanical properties (molecular weight increase). Disadvantages of the process variant described are the long preparation time and the disadvantageous properties previously mentioned.

[9024] DE 102004 005652.8 and DE 102004 005657.9 have previously proposed novel flow improvers for polyesters.

[0025] The prior art discloses additives for improving flow in polyamides:

[0026] FR-A 28 33 603; hyperbranched polyamides as rheology modifiers for PA

[0027] U.S. Pat. No. 6,541,599 soluble hyperbranched polyamides and their use, inter alia as rhoology modifiers for thermoplastics

[0028] DE-A 102 51 294: branched PA as additive for transparent, amorphous PAs

[0029] DE-A 102 55 044: terminal-polyfunctional (i.e. branched, hyperbranched, dendritic) polymers of the group of polyesters, polyglyterolls, polyethers as rheology modifiers in PA and PBT. (B089) EPA 682 OS7 describes an improved-flow prionwith a specific first generation 1.4-disministratural plepsyination dendrinor. DEA 19953 950 describes flowshie compounded polyamide materials with highly branched polyetheranides as reagent. The flow improver in this case first becomes agreement on addition of 10% of polyetheraniide. Universal applicability is lacking in both examples, and moreover the synthesis of the dendrinor described is inconvenient and expensive. The improvement in flow serivity, properties is also observed, and extend of the mechanical properties is also observed, and a first in mechanical

[0031] It was therefore an object of the present invention to provide thermoplastic polyamide molding compositions which have good flowability together with good mechanical properties.

[9032] Accordingly, the molding compositions defined at the outset have been found. Preferred embodiments are found in the subclaims.

[0033] The inventive molding compositions comprise, as component A), from 25 to 99% by weight, preferably from 30 to 98% by weight, and in particular from 40 to 95% by weight, of at least one polyamide.

[0034] The polyamides of the inventive molding compesitions generally have a viscosity number of from 90 to 350 ml/s, preferably from 110 to 240 ml/g, determined in a 0.5% strength by weight solution in 95% strength by weight sulfurie acid at 25° C. to 180 307.

[0035] Semicrystalline or amorphous resins with a molecular weight (weight-average) of at least 5000, e.g. those described in the American patent specifications U.S. Pat. Nos. 2,071,250, 2,071,251, 2,130,523, 2,130,948, 2,241,322, 2,312,966, 2,512,606 and 3,393,210 are preferred.

[0036] Examples of these are polyemides derived from lactans having from 7 to 13 ring members, e.g. polycaprolactam, polycaprylelactam, and polylaurolactam, and also polyamides obtained via reaction of dicarboxylic acids with diamnico.

[0037] Dicarboxylic acids which may be used are alkanedicarboxylic acids having from 6 to 12, in particular from 6 to 10, carbon atoms, and aromatic dicarboxylic acids. Acids which may be mentioned here merely as examples are adiptic acid, arolaic acid, sebacic acid, dodecanedioic acid and tereshibatic and/or isonbhalic acid.

[0038] Particularly suitable diamines are alkanediamines luving from 6 to 12, in particular from 6 to 8, carbon atoms, and also m-xylylenediamine, di(4-aminophenyl)methane, di(4-aminophenyl)propane, 2,2-di(4-aminopyleohexyl)propane or 1,5-diamino-2-methylenetiam

[0039] Preferred polyamides are polyhexamethyleneadipartide, polyhexamethylenesebeamide and polycaprolactam, and also yolon-6/6, for copolyamides, in particular having a proportion of from 5 to 95% by weight of caprolactam units.

[0040] Other suitable polyamides are obtainable from w-aminoalkyl nitriles, e.g. aminocapronitrile (PA 6) and adipodinitrile with hexamethylenedismine (PA 66) via what is known as direct polymerization in the presence of water, for example as described in DE-A 10313681, EP A 1198491 and EP 922065.

[0041] Mention may also be made of polyamides obtainable, by way of example, via condensation of 1,4-diaminobutane with delipie acid at an elevated temperature (nylon-4,6). Preparation processes for polyamides of this structure are described by way of example in EP-A 38 094, EP-A 38 582, and EP-A 39 524.

[0042] Other suitable examples are polyamides obtainable via copolymerization of two or more of the abovementioned monomers, and mixtures of two or more polyamides in any desired mixing ratio.

[0043] Other polyamides which have preven particularly advantageous are semiaromatic copolyamides, such as PA 66/G and PA 66/6T, where the triamine content of these is less than 0.5% by weight, preferably less than 0.3% by weight (see EPA 299 444 and EPA 667 367).

[0044] Preferred polyamides are composed of:

[0045] A1) from 20 to 90% by weight of units which derive from terephthalic acid and from hexamethylenedi-

[8046] A2) from 0 to 50% by weight of units which derive from e-caprolactam, and

[8047] A3) from 0 to 80% by weight of units which derive from adinic acid and from hexamethylenediemine.

[9048] A4) from 0 to 40% by weight of other polyamideforming monomers, where the proportion of component (A2) or (A3) or (A4) or their mixtures is at least 10% by weight.

[0049] Component A1) comprises from 20 to 90% by weight of units which derive from terephthalic acid and from became the length of the component of the compone

[0050] The copolyamides comprise, alongside the units which derive from terephthalic acid and from hexamethylenediamine, units which derive from c-caprolactam and/or units which derive from adjuc acid and from hexamethylenediamine and/or units which derive from other polyamideforming morphomers.

[0951] The proportion of units which derive from e-caprolactam is at most 50% by weight, professibly from 20 to 50% by weight, in particular from 25 to 40% by weight, in particular from 25 to 40% by weight, while the proportion of units which derive from adipic acid and from hexamethylenedificant is up to 80% by weight, preferably from 30 to 75% by weight, and in particular from 55 to 60% by weight.

[0052] The copolyamides may also comprise not only units from e-caprolactam but also units from adipin acid and hexamethyfenedizamine; in this case, it is advantageous for the proportion of units free from stronging groups to be at least 10% by weight, preferably at least 20% by weight. The ratio of the units which derive from e-caprolactam and from hexamethylenedizamine is not subject to

[0053] Polyamides having from 50 to 80% by weight, in particular from 60 to 75% by weight, of units which derive from terephthalic acid and from hexamethylemediamine (units A1)) and from 20 to 50% by weight, professbly from 25 to 40% by weight, of units which derive from ε-caprolectam (units A2)) have proven particularly advantageous for many applications.

[0054] The semiaromatic copolyamides can comprise, alongside the units Al) to A3) described above, amounts of up to 49% by weight, preferably from 10 to 30% by weight, and in particular from 20 to 30% by weight, of other polyamide-forming monomers A4), these being known from other polyamides.

[0057] The following constitutions of component (A) are particularly preferred here:

[0058] A1) from 65 to 85% by weight of units which derive from terephthelic acid and from hexamethylenedi-

[8059] A4) from 15 to 35% by weight of units which derive from isophthalic acid and from hexamethylenediamine, or

amine, or [0060] A1) from 50 to 70% by weight of units which derive from terepithalic acid and from hexamethylenedi-

[0061] A3) from 10 to 20% by weight of units which derive from adipic acid and from hexamethylenedismine,

omine and

[0062] A4) from 20 to 30% by weight of units which derive from isophthalic acid and from hexamethylenedi-

[0063] If compenent (A) comprises symmetrical dicarboxylic acids in which the earboxy groups are in para-8 position, it is advisable to structure these with (A) and (A2) or (A1) and (A2) as ternary copolyamides, since otherwise the ecopolyame would have an executively high melting point and would only melt with decomposition, this being undecirable.

[884] Other copolyamides which have proven particularly advantageous are those semigromatic copolyamides

whose triamine content is less than 0.5% by weight, preferably less than 0.3% by weight.

[9063] Semiaronatic copolyamides propared by most of the known processes (cf. U.S. Ph. No. 4,693,166) have triamine contents above 0.5% by weight, and this least to impairment of product quality and to problems in contents preparation. A particular triamine which may be mentioned of causing these problems is differentifying the intertional process.

[9066] Copolyamides with low triamine content have lower melt viscosities at the same solution viscosity, when compared with products of the same constitution which have high triamine content. This improves not only processibility but also product properties considerably.

[0067] The melting points of the semiaromatic copolyumides are in the range from 270 to 325° C., preferably from 280 to 310° C., and this high melting point is also associated with a high glass transition temperature which is generally above 75° C., in particular above 85° C. (in the dry state).

[9068] Binary copolyamides based on terephthalic acid, hexamethylenediamine and e-caprolactam have meliting points in the region of 300° C. and (in the dys state) a glass transition temperature above 110° C. if they have about 70% content of units which derive from terephthalic acid and from hexamethylenediamine.

[9069] Binary copolyamides based on terephinhile seid, adapte seid, and hexamethylene-diname achieve meiting points of 300° C. and above even at relatively low contents or about 55% by weight of units derived from terephinhile acid and hexamethylenediamine (HiMD), but the glass transition temperature here is not quite as high as for binary copolyamides which comprise c-caprolactam instead of adipte acid of the comprise com

[9070] According to the invention, the semiaromatic copolyamides are those whose degree of crystallinity is >10%, preferably >15%, and in particular >20%.

[9071] The degree of crystallinity is a measure of the fraction of crystalline fragments in the copolymer and is determined via X-ray diffraction.

[9072] The processes described in EP-A 129 195 and 129 196 can be used to prepare the preferred semiaromatic copolyamides with low triamine content.

[9073] It is, of course, also possible to use mixtures of the semiaromatic copolyamides, the mixing ratio here being as desired.

[0074] The following list, which is not comprehensive, comprises the polyamides A) mentioned and other polyamides A) for the purposes of the invention, and the monomers present:

An posymers	_
PA 4	Pyrrolidone
PA 6	e-Caprolactam
PA 7	Exhanolactum
PA 8	Caprylolactum
PA 9	9-Aminopelargonic sold
PA 11	11-Aminoundreanole acid
PA 12	Lauroisctam

AA/BB polyment:

PA 66

-continued Tetramethylenediamine, adipic acid Hexamethylenediamine, adipic acid

PA 69	Hexamethylenediamine, azetaje acad
PA 610	Hexamethylenediamine, schoole said
PA 612	Hexamethylenediamine, decanedicarboxylic seld
PA 613	Hexamethylenedismine, undecanedicarboxylic scie
PA 1212	1.12-Dodecanediamine, decanedicarboxylic acid
PA 1313	1.13-Disminotridecane, undecanedicarboxylic acid
PA 6T	Hexamethylenediamine, terephthalic acid
PA MXD6	m-Xylvienediamine, adipic sold
AA/BB nofymers:	
PA 6[Hexamethylenedismine, populthalic acid
PA 6-3-T	Trimethythesemethyleaedismine, terophthalic acid
PA 6/6T	(see PA 6 and PA 6T)
PA 6/66	(see PA 6 and PA 66)
PA 6/12	(see PA 6 and PA 12)
PA 66/6/610	(see PA 65, PA 6 and PA 610)
PA 6I/6T	(see PA 61 and PA 6T)
PA 6I/6T/PACMT	as PA 61/6T + diaminodicyclohexylmethane,
	terephthalic acid
PA 6T/6I/MACMT	as 63/6T + dimethyldiaminocyclohexylmethage,
	terephthalic acid
PA 6T/6I/MXDT	sa PA 61/6T + m-xylytenediamiae, terephthalic sci
PA 12/MACMI	Lauroinstein, dimethyldiaminodicyclohexylmethen
	isophthalic acid
PA 12/MACMT	Laurolactum, dimethyldiaminodicyclohexylmethan
	terephthalic acid
PA PDA-T	Phenylegedismine, terephthallo acid

[6975] The inventive molding compositions comprise, as component S), from 0.01 to 50% by weight, preferably from 0.5 to 20% by weight, and in particular from 0.7 to 10% by 0.5 to 20% by weight, and in particular from 0.7 to 10% by 0.5 to 20% by 0.5 to 20

[8076] For the purposes of this invention, hyperbranched polycorborates BJD are non-crossibled mineroundersules having hydroxy groups and carbonate groups, these having hotos structural and neclecular nomaliforation; Their structure may firstly be based on a central molecules in the same way as denderiment, but with nomaliform chain length of the branches. Secondly, they may also have a linear structure with functional predient groups, or else they may combine the two extremes, having linear and branched molecular proprients. See also F. J. Floys, J. Am. Chem. Soc. 152, 74, 2765. and 1. Frey et al., Chem. Bar. J. 2000, 6, no. 14, 2499 for the clinical cost of clientfaster and hyperbranched polymer.

[6077] "Hyperformacher" in the context of the present invention means that the degree of branching (DB), i.e. the average number of dendritic linkages plus the average number of edit group per molecule, is from 10 to 95.9%, preferably from 20 to 95%, particularly referably from 20 to 95%. "Dendritiesed" in the coetest of the present invention of the present the present the present invention of the present the present the present invention of the present the present the present invention of the definition of "degree of branching".

[0078] Component B1) preferably has a number-average molar mass M_n of from 100 to 15 000 g/mol, preferably from 200 to 12 000 g/mol, and in particular from 500 to 10 000 g/mol (GPC, PMMA standard).

[0079] The glass transition temperature Tg is in particular from -80 to +140° C., preferably from -60 to 120° C. (according to DSC. DIN 53765).

[0080] In particular, the viscosity (mPas) at 23° C. (to DIN 53019) is from 50 to 200 000, in particular from 100 to 150 000, and very particularly preferably from 200 to 100 000.

[0081] Component B1) is preferably obtainable via a process which comprises at least the following steps:

[9882] a) neution of at least one organic carbonate (A) of the gomenl formula RO((COO)[4], with at least one aliphatic, aliphatic/aromatic, or aromatic alcohol (B) which has at least 30H groups, with elimination of alechols ROH to give one or more condensates (S), where each R, independing of the others, is a striight-chain or branched aliphatic, aromate/daliphatic, or aromatic hydrowhere the radicals R may also have bonding to one nonder to form a ring, and n is a whole number from 1 to 5, or

[0083] ab) reaction of phosgene, diphosgene, or triphosgene with abovementioned alcohol (B) with elimination of hydrogen chloride and

[0084] b) intermolecular reaction of the condensates (K) to give a high-functionality, highly branched, or high-functionality, hyperbranched polycarbonate,

[9085] where the quantitative proportion of the OH groups to the carbonates in the resortion mixture is selected in such a way that the condensates (K) have an average of cither one carbonate group and more than one OH group or one OH group and more than one carbonate group.

[9086] Starting materials which may be used comprise phosphene, diphosgene, or triphosgene, preference being given to organic carbonates.

[6697] Back of the neticeals is of the organic enchosism (A) used as sturing, mental and having the general formals RO(CO/ORS is independently of the others, a straight-chain or beneched injustic, contractival plantin, contractival plantin, or anomalo hydrocarbon medical having from 1 to 20 carbon arouns. The two nedicals Rum yale have benefiting to not notice 1st for an ring. The residual is preferably an aliphatic hydrocarbon miderial, and particularly preferably a nitrigate that on branched alley mideral baving from 1 to 5 carbon arouns, or banched alley mideral baving from 2 to 5 carbon arouns, or anothering on a machinated professional profes

[0088] Use is particularly made of simple carbonates of the formula RO(CO)OR; n is preferably from 1 to 3, in particular 1.

[0889] By way of example, dialkyl or dianyl corbonates may be prepared from the reaction of nilpatis, annihatric, or amonatic alcohola, preferably monoslocholos, with posgene. They may also be prepared by way of oxidarive carbonylation of the alcoholo or phenois by means of CO in the presence of rache metals, oxygen, or NO, In relation to preparation methods for diarryl or dialkyl carbonates, see edition, 2000 [Sections Relocates, Vedena Wijle, VVCH, decition, 2000 [Sections Relocates, Vedena Wijle, VVCH, [6999] Estraples of saitable curbanates comprise aliphoris, accurate l'alphalier or sementier conbouete, ende se ethylene curbonete, propylene 1,2- or 1,3-curbonate, displasive proposition de l'accurate de l'accurate de l'accurate des displasments de l'accurate de l'accurate de l'accurate de l'accurate distribution de l'accurate de l'accurate de l'accurate de l'accurate de distribution de l'accurate d

[0091] Examples of carbonates in which n is greater than 1 comprise dialkyl dicarbonates, such as di(tert-butyl)dicarbonate, or dialkyl tricarbonates, such as di(tert-butyl)tricar-

[0992] It is preferable to use aliphatic carbonates, in particular those in which the radicals comprise from 1 to 5 carbon atoms, e.g. dimethyl carbonate, diethyl carbonate, dipropyl carbonate, dibutyl carbonate, or diisobutyl carbonate.

[9093] The organic carbonates are reacted with at least one aliphatic alcohol (B) which has at least 3 OH groups, or with mixtures of two or more different alcohols.

[0094] Examples of compounds having at least three OH groups comprise glycerol, trimethylolmethane, trimethylolethane, trimethylolpropane, 1,2,4-butanetriol, tris(hydroxymethyl)amine, tris(hydroxyethyl)amine, tris(hydroxypropyl)amine, pentserythritol, diglycerol, triglycerol, polyglycerols, bis(trimethylolpropane), tris(hydroxymethyl-)isocyanurate, tris(hydroxyethyl)isocyanurate, phloroglucinol, trihydroxytoluene, trihydroxydimethylbenzene, phloroglucides, hexahydroxybenzene, 1.3.5-benzenetrimethanol. 1,1,1-tris(4'-hydroxyphenyl)methane, 1,1,1-tris(4'-hydroxyphenyl)ethane, bis(trimethylolpropane), or sugars, e.g. glucose, trihydric or higher-functionality polyetherols based on trihydric or higher-functionality alcohols and ethylene oxide, propylene oxide, or butylene oxide, or polyesterols. Particular preference is given here to glycerol, trimethylolethane, trimethylolpropane, 1,2,4-butanetriol, pentserythritol, and their polyetherols based on ethylene oxide or propylene oxide

[9095] These polyhydric alcohols may also be used in a mixture with dihydric alcohols (B'), with the proviso that the average OH functionality of the totality of all of the alcohols used is greater than 2. Examples of suitable compounds having two OH groups comprise ethylene glycol, diethylene glycol, triethylene glycol, 1,2- and 1,3-propanediol, dipropylene glycol, tripropylene glycol, neopentyl glycol, 1,2-, 1,3-, and 1,4-butancdiol, 1,2-, 1,3-, and 1,5-pentanediol, hexanediol, cyclopentanediol, cyclohexanediol, cyclohexanediol, cyclohexanediol, cyclohexanediol, cyclohexanediol, cyclohexanediol, cyclohexanediol, cyclohexanediol anedimethanol, bis(4-hydroxycyclohexyl)methane, bis(4-hydroxycyclo-hexyl)ethane, 2,2-bis(4-hydroxycyclohexyl-1,1'-bis(4-hydroxyphenyl)-3,3,5trimethyleyelohexane, resorcinol, hydroquinone, dihydroxyphenyl, bis(4-bis(hydroxyphenyl)sulfide, bis(4hydroxyphenyl)sulfone, bis(hydroxymethyl)benzene, bis(hydroxymethyl)toluene, bis(p-hydroxyphenyl)methane, bis(p-hydroxyphenyl)ethane, 2,2-bis(p-hydroxyphenyl)pro-pane, 1,1-bis(p-hydroxyphenyl)cyclohexane, dihydroxybenzophenone, dihydric polyether polyels based on ethylene oxide, propylene oxide, butylene oxide, or their mixtures, polytetrahydrofuran, polycaprolactone, or polyesterols based on diols and diearboxylie acids. [9094] The diols serve for fine adjustment of the properties or for the polyambons. Utrus is made of Highirds selectors, the notion of distyricis actions, the notion of distyricis actions. If the notion of distyricis actions are the notion of distyricis actions are the notion of the properties of the polyambons. The amount of the desired properties of the polyambons. The amount of the schools (3P) give generally from 0 to 39 and 9%, based on the centre amount of the totality of all of the alcohols (3P) and (9P). The amount is professibly from 0 to 25 and 5%, and very particularly prefensibly from 0 to 25 and 5%, and 25 and

[0097] The reaction of phosgene, diphosgene, or triphosgene with the alcohol or alcohol mixture generally take place with elimination of hydrogen chloride, and the reaction of the carbonates with the alcohol or alcohol mixture to give the inventive high-functionality highly branched polycarbonate takes place with elimination of the monohydric alcohol or phenol from the carbonate molecule.

[6998] After the maction, i.e. without further modification, the high-functionality highly branched polycurbonates formed by the invention process have termination by plotony groups and/or by entroline process. have termination by plotony groups and/or by entroline groups. They have good establiship in various solvents, e.g. in worter, election, such as mediane, eleman, but intunal, elemboly such entralises, extension, etc. 2-buttanens, etc. 3 section, further mattern, architecture, architecture, methodolyment, dimethylem-manide, dimethylenotamides. N-methylypronidone, efciyl-sensible, dimethylenotamides. N-methylypronidone, efciyl-sensible, dimethylenotamides.

[9999] For the purposes of this invention, a high-functionnial propherations is a product which, is foliated the entroluse groups which form the polymer felerion, further has at least three, preferrably a least its, more perfermily at least its, terminal or penduct functional groups. The functional groups are contained groups and/or to figures. There is ingroups are contained groups and the figures. There is intermediated three properties, the properties of the properties, and the properties, and the properties, such as high viscosity or poor subshilty. The highfunctionality polyse-absents of the prevent invention month, properties, and the properties of the prevention of the product groups, research as 200 terminal or product functional groups, research as the product function of product functional propage.

[0109] When preparing the high-functionality polycarbonness B1), it is necessary to adjust the ratio of the compounds as B1), it is necessary to adjust the ratio of the compounds when the property of the compound of the compound of the way that the implement resultant condensate (bereinfler tensed condensate (KI) comprises an average of either one candonate group or entamony group and more than one of the group or one Oil group and more than one carbonate group or entamony group. The implement structure of the condensate (K) composed of a carbonate (A) and a di- or polyulcolo (II) have reasons in the mrangement X₁, or Y, x, where X is a carbonate group, Y is a hydroxy group, and n is generally a number from 1 to 6, preferredly from 1 to 4, particularly preferredly from 1 to 5. The reserve group which is the property of the compound of the property of the composition of the composition of the composition of the composition of the particularly preferred by from 1 to 5. The reserve group which is the property of the composition of the composition of the composition of the particularly preferred by from 1 to 5. The reserve group which is a property of the composition o

[0101] By way of example, if during the preparation of the simplest condensate (K) from a carbonate and a dihydric alcohol the reaction ratio is 1:1, the average result is a molecule of XY type, libratrated by the general formula 1.

[0102] During the preparation of the condensate (K) from a carbonate and a trihydric alcohol with a reaction ratio of 1:1, the average result is a molecule of XY₂ type, illustrated by the general formula 2. A carbonate group is focal group here.

[0103] During the preparation of the condensate (K) from a curbonate and a tetrahydric alcohol, likewise with the reaction ratio 1:1, the average result is a molecule of XY₃ type, illustrated by the general formula 3. A carbonate group is focal group here.

[0104] R in the formulae 1-3 has the definition given at the outset, and R¹ is an aliphatic or aromatic radical.

[0105] The condensate (K) may, by way of example, also be prepared from a carbonate and a trihydric alcohol, as illustrated by the general formula 4, the moder reaction ratio being 2:1. Here, the average result is a molecule of X₂Y type, an OH group being focal group here. In formula 4, R and R¹ are as defined in formulae 1:3.

[0106] If difunctional compounds, e.g. a dicarbonste or a diol, are also added to the components, this extends the chains, as illustrated by way of example in the general formula 5. The average result is again a molecule of XY, type, a carbonate cruture being focal ground.

[0107] In formula 5, R^a is an organic, preferably aliphatic radical, and R and R^a are as defined above.

[9188] It is also possible to use two or more condensates (6) for the synthesis, Firstly, two or more solvabile and, respectively, two or more corbonates may be used here. Furthermore, mixtures of various condensates of different structures can be obtained via the selection of the ratio of the activations and, respectively, the phospson. This will be illustrated taking the example of the procession of a corbonate with a tribuplier alterbal. If the expectation of a corbonate with a tribuplier alterbal. If the corbonate with a tribuplier alterbal, If the corbonate with a tribuplier alterbal. If the corbonate with a tribuplier alterbal, If the corbonate with a tribuplier alterbal. If the corbonate with a tribuplier alterbal, If the corbonate with a tribuplier alterbal. If the corbonate with a tribuplier alterbal, If the corbonate with a tribuplier alterbal is the corbonate with a tribuplier alterbal. If the corbonate with a tribuplier alterbal is a tribuplier and 2:1 the product is an attitute of XY, and XY, productles.

[909] According to the invention, the simple condensates (SC) described by swe of trample in the formulae 1-5 perfectually react internalized by form high-function-simply produced by the section of the produced by the prod

[0110] In one preferred embodiment, the condensation reaction is carried out in bulk. The phanol or the monohydric

alcohol ROH liberated during the reaction can be removed by distillation from the reaction equilibrium to accelerate the reaction, if appropriate at reduced pressure.

[0111] If removal by distillation is intended, it is generally advisable to use those carbonates which liberate alcohols ROH with a boiling point below 140° C, during the reaction.

[6112] Carlalysts or casalyst mixtures may also be added to accelerate the receion. Solitable carlibrs are compounds which enalyze esterification or transesterification reactions, e.g. astall meat layloractions, shall used corbusous, shall meat layloraction, surface and compounds, or portains, or of cessins, unctured smaller, surface situation, or of cessins, unctured smaller, organociation, compounds, phasphantium compounds, organociationium, organociation, are organication, organociation, and compounds of the compound of the compounds of the

[9113] It is preferable to use potassima hydroxide, potassium carbonate, potassium hydrogeneurbonate, disabievichoctune (DABCO), diszabievy-lononene (DBN), diszabicy-lonodenene (DBU), imidzones, suot as imicken, lamethy-limidzozle, or 1,2-dimethy-limidzozle, titarium tetratuboxide, titarium tetrasporpovolcie, dibutylim di dibutylim dilaurute, stanonous dioctoste, zirconium scetylacetonate, or mixtures thereol.

[0114] The amount of catalyst generally added is from 50 to 10 000 ppm by weight, preferably from 100 to 5000 ppm by weight, based on the amount of the alcohol mixture or alcohol ward.

[0115] It is also possible to control the intermolecular polycondensation reaction via addition of the suitable canityst or else via selection of a suitable temperature. The average molecular weight of the polymer (P) may moreover be adjusted by way of the composition of the starting components and by way of the residence time.

[0116] The condensates (K) and the polycondensates (P) prepared at an elevated temperature are usually stable at room temperature for a relatively long period.

[0117] The nature of the condensates (K) permits polycondensates (F) with different structures to result from the condensation reaction, these having branching but no rossilation, Purtlemence, in the ideal case, the polycondensates (P) have either one carbonate group as focal group and more than two CH groups or elso use OH group as focal group and more than two carbonate groups. The number of condensates (E) have been only the condensation of the condensate (E) have and aft the exerce of polycondensation.

[0118] By way of example, a condensate (K) according to the general formula 2 can react via triple intermolecular condensation to give two different polycondensates (P), represented in the general formulae 6 and 7.

[0119] In formula 6 and 7, R and R1 are as defined above.

[0120] There are various ways of terminating the intermolecular polycondensation reaction. By way of example, the temperature may be lowered to a range where the reaction stops and the product (K) or the polycondensate (P) is storace-stable.

[9121] It is also possible to deactivate the catalyst, for example in the case of basic catalysts via addition of Lewis acids or protonic acids.

[8122] In mother embodiment, as soon as the intermolecular reaction of the condensate (G.) has produced a polycondensate (P) with the desired degree of polycondensation, a product having groups reactive lowered the foal reaction. For example, in the case of a carbonate group as floral group, by way of example, a moso, dl. or polyamien may be added. In the case of a bydroxy group as fixed group, by the way of example, a moso, dl. or polyamien to be added to the case of a bydroxy group as fixed group, by way of example, a moso, dl. or polyamien of the control of the product of the product of the production of the product of

[0123] The inventive high-functionality polycarbonates are mostly prepared in the pressure range from 0.1 mbar to 20 bar, preferably at from 1 mbar to 5 bar, in reactors or reactor caseades which are operated batchwise, semicontinuously, or entitionally, or entitionally.

[0124] The inventive products can be further processed without further purification after their preparation by virtue of the abovementioned adjustment of the reaction conditions and, if appropriate, by virtue of the selection of the suitable

[0125] In another preferred embodiment, the product is stripped, i.e. freed from low-molecular-weight, volatile compounds. For this, once the desired degree of conversion has been achieved the catalyst can optionally be descrivated and the low-molecular-weight volatile constituents, e.g. montalcoloids, phenols, carbonates, hydrogen chloride, or high-volatility oligomers or cyclic compounds can be removed by distillation, if appropriate with introduction of a pas, preferably nitrogen, carbon dioxide, or air, if appropriate a treduced pressure.

[0126] In another preferred embodiment, the inventive polycarbonates may acquire other functional groups besides the functional groups equired by virtue of the reaction. The functionalization may take place during the process to increase molecular weight, or else subsequently, i.e. after completion of the actual polycondensation.

[0127] If, prior to or during the process to increase molecular weight, components are added which have other functional groups or functional elements besides hydroxy or carbonate groups, the result is a polycarbonate polymer with randomly distributed functionalities other than the carbonate or hydroxy groups.

[9128] Effects of this type can, by way of example, be achieved via addition, during the polycondensation, of compounds which bear other functional groups or functional elements, such as mercapto groups, primary, secondary or tertiary amino groups, ether groups, derivatives of carboxy-lic acids, derivatives of sulfonic acids, derivatives of phosphonic acids, silane groups, siloxane groups, arvi radicals, or long-chain alkyl radicals, besides hydroxy groups, carbonate groups or carbamoyl groups. Examples of compounds which may be used for modification by means of carbamate groups are ethanolamine, propanolamine, isopropanolamine, 2-(bu-tylamino)ethanol, 2-(cyclohexylamino)ethanol, 2-amino-1butanol, 2-(2-aminoethoxy)ethanol or higher alkoxylation products of ammonia, 4-hydroxypiperidine, 1-hydroxyethylpiperazine, diethanolamine, dipropanolamine, diisopropanolamine, tris(hydroxymethyl)-aminomethane, tris(hydroxyethyl)aminomethane, ethylenediamine. propylenediamine, hexamethylenediamine or isophoronedi-

[9129] An example of a compound which can be used for modification with mercepts groups is merceptechnol. By way of example, tertiary amine groups can be produced via incorporation of Nenethyleidenhooltemine. By way of example, their gaves may be generated via occondensation of disputio or polytydrid polyethenols. Long-chain ally radicals can be introduced via reaction with lange-chain ally radicals can be introduced via reaction with lange-chain alloracidist, and reaction with ally or any dissocyamates ground polytyme and having ally 1, 91, and trethous

[0130] Addition of dicarboxylic acids or tricarboxylic acids, or, for example, dimethyl terephthalate, or tricarboxylic esters can produce ester groups.

[0131] Subsequent functionalization can be achieved by using an additional step of the process (step c)) to react the resultant high-functionality highly branched, or high-functionality hyperbrunched polycarbonate with a suitable functionalizing reagent which can west with the OH and/or carbonate groups or carbonary groups of the polycarbonate.

[0132] By way of example, high-functionality highly branched, or high-functionality hyperbranched polycarbonates comprising hydroxy groups can be modified via addition of molecules comprising acid groups or comprising isocyanate groups. By way of example, polycurbonates comprising acid groups can be obtained via reaction with compounds comprising anhydride groups.

[0133] High-functionality polycarbonates comprising hydroxy groups may moreover also be converted into high functionality polycarbonate polyether polyols via reaction with alkylene oxides, e.g. ethylene oxide, propylene oxide, or butylene oxide.

[0134] A great advantage of the process is its cost-offictiveness. Both the reaction to give a condensate (K) or polycondensate (P) and also the reaction of (K) or (P) to give polycarbonates with other functional groups or elements can take place in one reactor, this beling advantageous technically and in terms of cost-effectiveness,

[0135] The inventive molding compositions may comprise, as component B2), at least one hyperbranched polyester of AxBy type, where

x is at least 1.1, preferably at least 1.3, in particular at least 2

y is at least 2.1, preferably at least 2.5, in particular at least

[0136] Use may also be made of mixtures as units A and/or

B. of course

[0137] An A_xB_y type polyester is a condensate composed of an x-fractional molecule A and a y-functional molecule B. By way of example, mention may be made of a polyester composed of adipic acid as molecule A (x=2) and glycerol as molecule B (y=3).

[9138] For the purposes of this invention, hypotheranched opposters B23 are noncrevilinked merconfectolate having both amount of the policy of the property of the policy of the policy

[9139] "Hyperbranched" in the context of the present invention means that the degree of branching (DB), i.e. the average number of dendritic linkages plus the average number of end groups per molecule, is from 10 to 99.9%, perferably from 20 to 99%, particularly preferably from 20 to 95%.

[0140] "Dendrimeric" in the context of the present invention means that the degree of branching is from 99 9 to 100%. See H. Frey et al., Acta Polym. 1997, 48, 30 for the definition of "degree of branching".

[0141] Component B2) preferably has an M_n of from 300 to 30 000 g/mol, in particular from 400 to 25 000 g/mol, and very particularly from 500 to 20 000 g/mol, determined by means of GPC, PMMA standard, dimethylacetamide eluent

[0142] B2) preferably has an OH number of from 0 to 600 mg KOH/g of polyecter, preferably of from 1 to 500 mg

KOH/g of polyester, in particular from 20 to 500 mg KOH/g of polyemer to DIN 53240, and proferably a COCH number of from 0 to 600 mg KOH/g of polyester, preferably from 1 to 500 mg KOH/g of polyester, and in particular from 2 to 500 mg KOH/g of polyester,

[0143] The T_g is prefembly from -50° C. to 140° C_γ and in particular from -50 to 100° C. (by means of DSC, to DIN 53765).

[0144] Preference is particularly given to those components B2) in which at least one OH or COOH number is greater than 0, preferably greater than 0.1, and in particular greater than 0.5.

[0145] The inventive component B2) is in particular obtainable via the processes described below, inter alia by reacting

[0146] (a) one or more dicarboxylic acids or one or more derivatives of the same with one or more at least trihydric alcohols

'n

[0147] (b) one or more tricerboxylic acids or higher polycerboxylic acids or one or more derivatives of the same with one or more diols

in the presence of a solvent and optionally in the presence of an inorganic, organometallic, or low-molecular-weight organic catalyst, or of an enzyme. The reaction in solvent is the preferred preparation method.

[0148] For the purposes of the present invention, highfunctionality hyperbranched polyesters B2) have molecular and structural nonuntomity. Their molecular nonunitormity distinguishes them from dendrimers, and they can therefore be prepared at considerably lower cost.

[949] Azoneg tine disenboxylic neide which can be reacted according to varies (a) sur, by way of compile, oxalia exid, matonia acid, succinic neid, gutaria neid, specia exid, subcinic acid, specia exid, specia exid, specia exid, special exid, special exid, special exid, special exid, special exid, special exid, exide activa exid, exide activa expolherates 1-3, disenboxylic acid, exid activa expolherates 1-3, disenboxylic acid, exid and trans-cycloherates 1-4, disenboxylic acid, exid and trans-cycloherates 1-4, disenboxylic acid, exid and trans-cyclopentare 1-3, disenboxylic acid, and exis and trans-cyclopentare 1-3, disenboxylic acid, exid and trans-cyclopentare 1-3, disenboxylic acid, and exis and trans-cyclopentare 1-3, and trans-cyclopentare 1-3, and trans-cyclopentare 1-3, and trans-cyclopentare 1-3

[0150] and the abovementioned dicarboxylic acids may have substitution by one or more radicals selected from

[9151] C₁-C_{1,0}-allyl groups, such as methyl, ethyl, n-proryl, isopropyl, a buryl, isobetyl, see buryl, set buryl, a prolyl, isopentyl, see-pentyl, neupentyl, 1,2-dimethylprupyl, isosamyl, n-hexyl, isohexyl, see-hexyl, n-heptyl, isoheptyl, n-o-tyl, 2-ethylhexyl, n-nonyl, and n-decyl,

[0152] C₃ C₁₂ cycloalkyl groups, such as cyclopropyl, cyclobatyl, cycloactyl, cyclohexyl, cyclohexyl, cyclonopyl, cycloaceyl, cyclonacyl, and cyclodocyl, cyclonecyl, and cyclohexyl, and cyclohexyl, and cyclohexyl, and cyclohexyl, and cyclohexyl,

[0153] alkylene groups, such as methylene or ethylidene, or

[0154] C_6 - C_{14} -aryl groups, such as phenyl, 1-naphthyl, 2-naphthyl, 1-anthryl, 2-anthryl, 9-anthryl, 1-phenanthryl,

2-phenanthryl, 3-phenanthryl, 4-phenanthryl, and 9-phenanthryl, preferably phenyl, 1-naphthyl, and 2-naphthyl, perticularly preferably phenyl.

[9155] Examples which may be mentioned of representatives of substituted dicarboxylic acids are: 2-methylmalonic acid, 2-ethylmalonic acid, 2-phenylmalonic acid, 2-methylsuccinic acid, 2-ethylsuccinic acid, 2-phenylsuccinic acid, itsconic acid, 3-3-dimethyletatric acid,

[0156] Among the dicarboxylic acids which can be reacted according to variant (a) are also ethylenically unsaturated acids, such as maleic acid and funaric acid, and aromatic dicarboxylic acids, such as pluthatic acid, isophthalic acid or terephthalic acid.

[0.157] It is also possible to use mixtures of two or more of the abovementioned representative compounds.

[0158] The dicarboxylic acids may either be used as they stand or be used in the form of derivatives.

[0159] Derivatives are preferably

[0160] the relevant anhydrides in monomeric or else polymeric form,

[0161] mono- or dialkyl esters, preferably mono- or dimethyl esters, or the corresponding mono- or diethyl esters, or else the mono- and dialkyl esters derived from higher alcohols, such as n-propanol, isopropanol, n-butanol, isobutanol, ter-butanol, n-bertanol, n-bexanol,

[0162] and also mono- and divinyl esters, and

[0163] mixed esters, preferably methyl ethyl esters.

[0164] In the preferred preparation process it is also possible to use a mixture composed of a dicarboxylic acid and one or more of its derivatives. Equally, it is possible to use a mixture of two or more different derivatives of one or more dicarboxylic acids.

[0165] It is particularly preferable to use succinic acid, glutane acid, acipic acid, plutane acid, scophthalic acid, or the mono- or dimethyl ester thereof. It is very particularly preferable to use adipic acid.

[9164] Examples of at least trihydric alexhols witch may be reacted are glycorol, bitates 1,24-fix), apertance 1,35-triol, a-pectance 1,35-triol, a-became-1,25-triol, a-became-1,25-triol, a-became-1,35-triol, a-became-1

[0167] Examples of tricarboxylic acids or polycarboxylic acids which can be reacted according to variant (b) are benzene-1,2,4-tricarboxylic acid, benzene-1,3,5-tetracarboxylic acid, and melliticated

[0168] Tricarboxylic acids or polycarboxylic acids may be used in the inventive reaction either as they stand or else in the form of derivatives.

[0169] Derivatives are preferably

[0170] the relevant anhydrides in monomeric or else polymeric form,

[0171] mono-, di-, or trialkyl esters, preferably mono-, di-, or triankyl esters, or the corresponding mono-, di-, or triethyl esters, or else the mono-, di-, and triesters derived from higher alcohols, such as n-propanol, iso-propanol, n-butanol, isobutanol, retri-butanol, n-penanol, n-butanol, or else mono-, di-, or trivinyl esters

[0172] and mixed methyl ethyl esters.

[0173] For the purposes of the present invention, it is also possible to use a mixture composed of a tri- or polycarboxylic acid and one or more of its derivatives. For the purposes of the present invention it is likewise possible to use a mixture of two or more different derivatives after one or more tri- or polycarboxylic acids, in order to obtain component [32].

[0174] Examples of diols used for variant (b) of the present invention are ethylene glycol, propane-1,2-diol, propane-1,3-diol, butane-1,2-diol, butane-1,3-diol, butane-1,4diol, butane-2,3 diol, pentane 1,2 diol, pentane 1,3 diol, pentane-1,4-diol, pentane-1,5-diol, pentane-2,3-diol, pentane-2,4-diol, hexane-1,2-diol, hexane-1,3-diol, hexane-1,4diol, hexane-1,5-diol, hexane-1,6-diol, hexane-2,5-diol, heptane-1,2-diol, 1,7-heptanediol, 1,8-octanediol, 1,2-octanediol, 1,9-nonanediol, 1,10-decanediol, 1,2-decanediol, 1,12-dodecanediol, 1,2-dodecanediol, 1,5-hexadiene-3,4diol, cyclopentanediols, cyclohexanediols, inositol and derivatives, (2)-methylpentane-2,4-diol, 2,4-dimethylpentane-2,4-diol, 2-ethylhexane-1,3-diol, 2,5-dimethylhexane-2,5-diol, 2,2,4-trimethylpentane-1,3-diol, pinacol, diethyl ene glycol, triethylene glycol, dipropylene glycol, tripropylene glycol, polyethylene glycols HO(CH₂CH₂O)_n—H or polypropylene glycols HO(CH [CH₃]CH₂O)_n—H or mixtures of two or more representative compounds of the above compounds, where n is a whole number and n=from 4 to 25. One, or else both, hydroxy groups here in the abovementioned diols may also be substituted by SII groups. Preference is given to ethylene glycol, propane-1,2-diol, and diethylene glycol, triethylene glycol, dipropylene glycol, and tripropylene glycol.

[0175] The molar ratio of the molecules A to molecules B in the A_xB_y polyester in the variants (a) and (b) is from 4:1 to 1:4, in particular from 2:1 to 1:2.

[9.176] The sit least tribydric alcolols reacted according to variant (a) of the process may have hydroxy groups of which all have identical reactivity. Preference is also given here to at least tribydric alcohols whose OH groups initially have identical reactivity, but where reaction with a least one acid group can induce a fall-off in exactivity of the remaining OH groups as a result of state or electronic effects. By way of the action of the control of the control

[0177] However, the at least trihydric alcohols reacted according to variant (a) may also have hydroxy groups having at least two different chemical reactivities.

[0178] The different reactivity of the functional groups here may either derive from chemical causes (e.g. primary/ secondary/tertiary OH group) or from steric causes. [0179] By way of example, the triol may comprise a triol which has primary and secondary hydroxy groups, proferred example being glycerol.

[0180] When the inventive reaction is carried out according to variant (a), it is preferable to operate in the absence of diots and monohydric alcohols.

[9181] When the inventive reaction is carried out according to variant (b), it is preferable to operate in the absence of mono- or dicarboxylic acids.

[9182] The inventive process is carried out in the presence of a solvent. Beampies of satished compounds me hydro-curbons, such as passifies or armanise. Particularly smithole parafillar not n-alpeane and cyclothease. Particularly smithole parafillar such n-alpeane and cyclothease. Particularly scales able serounties are toluren, orthosylten, meta-sylten, para-sylten, xylene in the form of an isome matture, etyliphenzese, chiorehermen and ortho- and meta-dichlorohermen. Other very particularly satished solvents in the absence of four very particularly satished solvents in the absence of firme, and testones, such as methyl ethyl ketone and methyl isobard ketones.

[9182] According to the invention, the amount of polyunisided is at least of 16 by weight, heard on the weight of the testiring materials used and to be resired, preferrably at least 15 by weight, and particularly preferrably at least 16% by weight, it is also possible to use excesses of solvent, based on the weight of the starting materials used and to be reasoid, e.g. from 1.01 to 10 times the amount. Solvent amounts of more than 100 times the weight of the starting materials used more than 100 times the weight of the starting materials used rate veduces merkedly at materialy lower concentrations of the resectant, giving unconsumically 100 greation times.

[8184] To carry out the process preferred according to the invention, openions any be carried out in the presume of a dehydrating agent as additive, added at the start of the reaction. Skilable camples are molecular tieves, it perimies har 4. An molecular sieve, h. godge, and Na, SO, During the reaction it is also possible to add Utrate dedyvening agent or to replace dehydrating agent to the contract of the con

[9185] The process may be carried out in the absence of actidic actalysts. It is preferable to operate in the presence of an actidic inorganic, organometallic, or organic catalyst, or a mixture composed of two or more actidic inorganic, organometallic, or organic catalysts.

[9186] For the purposes of the present invention, examples of cacific inanguine cathysts are stilline airly, hospather early, a complex order, and a cardial companies and a prophosphore as self, a hautisma statis before, a fam, cacific silice, a [161-4]; in particular—5, and a solide stuntium motific. Examples or dust compounds which can be need as actific integrate catalyst are administra compounds of the general formula Al(OR), and tituates of the general formula fal(OR), where each of the naticular R may be identical or different and is selected independently of the others from.

[9187] C₁-C₁₀-alkyl radicals, such as methyl, ethyl, n-propyl, isoponyl, n-butyl, isobutyl, sec-butyl, ten-butyl, n-pentyl, isopontyl, sec-pontyl, noopontyl, 1,2-dimothylpropyl, isoamyl, n-hexyl, isohexyl, sec-hexyl, n-heptyl, isoheptyl, n-notyl, 2-ethylhexyl, n-nonyl, and n-decyl,

[0188] C₃-C_{1,2}-cycloalkyl mdicałs, such as cycloprupyl, cyclobutyl, cyclopenyl, cyclohoxyl, cyclohoxyl, cyclopenyl, tyl, cyclonanyl, cyclodecyl, cycloundecyl, and cyclodocecyl; preference is given to cyclopentyl, cyclobexyl, and cyclohetyl.

[0189] Each of the radicals R in Al(OR)₃ or Ti(OR)₄ is preferably identical and selected from isopropyl or 2-ethylboxyl.

[0190] Examples of preferred acidic organomentalic catalysts are selected from disklytin oxidos R_cSnO, where R is defined as above. A particularly preferred representative compound for acidic organometallic catalysts is di-a butyl-tin oxide, which is commercially available as "oxo-tin", or di-a-butylin dilaurate.

[8191] Preferred acidic organic catalysts are acidic organic compounds having, by way of example, phosphate group, sulfonic acid groups, militate groups, or phosphonic soids groups. Particular perfetence is given to sulfanic acids, such as para-tolume-sulfonic acid. Acidic ion exchangers may also be used as acidic organic catalysts, exp. polytical and the properties of the properties of the properties of the boat 2 and 3 of divinythermous.

[6192] It is also possible to use combinations of two or more of the abovementioned catalysts. It is also possible to use an immobilized form of those organic or organometallic, or else inorganic catalysts which take the form of discrete molecules.

[0193] If the intention is to use acidic inorganic, organometallic, or organic catalysts, according to the invention the amount used is from 0.1 to 10% by weight, preferably from 0.2 to 2% by weight, of catalyst.

[0194] The inventive process is carried out under inert gas, e.g. under carbon dioxide, nitrogen, or a noble gas, among which mention may particularly be made of argon.

[0195] The inventive process is carried out at temperatures of from 60 to 200° C. It is preferable to operate at temperatures of from 130 to 180° C., in particular up to 150° C., or below that temperature. Maximum temperatures up to 145° C. are particularly preferred, and temperatures up to 135° C. are very particularly preferred.

[6196] The pressure conditions for the inventive process are not critical per see. It is possible to operate at markedly reduced pressure, e.g., at from 10 to 500 mbar. The inventive process may also be carried out at pressures above 500 mbar. A resertion at timospheric pressure is preferred for reasons of simplicity; however, conduct at slightly increased pressure is simplicity; however, conduct at slightly increased pressure is operation at markedly increased pressure, e.g., at pressures upoperation at markedly increased pressure, e.g., at pressures in the 10 lbs IP Reservice on at simospheric pressure is preferred.

[0197] The reaction time for the inventive process is usually from 10 minutes to 25 hours, preferably from 30 minutes to 10 hours, and particularly preferably from one to 8 hours.

[0198] Once the reaction has ended, the high-functionality hyperbranched polyesters can easily be isolated, e.g. by removing the catalyst by filtration and concentrating the mixture, the concentration process here usually being cerried out at reduced pressure. Other work-up methods with good suitability are precipitation after addition of water, followed by washing and drying.

[0199] Component B2) can also be prepared in the presence of enzymes or decomposition products of enzymes (according to DB-A 101 63163). For the purposes of the present invention, the term acidic organic catalysts does not include the dicarboxylic acids reacted according to the invention.

(2000) It is perfonible to use liques or esterness. Lipues, and enterness with good estitability or condid cylinhorous, candid eylinhorous, Candida lipolytica, Candida ergoa, Candida stitus-Cons. Candida tili, Chromobacterim viscosum, Goolrichum candidum, Muco jewaticus, Muco midel, pig pancers, peredomonas epoca, p

[9261] The enzyme is prefembly used in immobilized form, for example on silica gel or Lewatif.9 Processes for immobilizing enzymes are known per se, e.g. from Kurt Faber, "Biotransformations in organic chemistry", 3rd edition 1907, Springer Verlag, Chippler 3.2 "Immobilization" pp. 345-336. Immobilized enzymes are commercially available, for example from Novoymes Biotich Inc., Demmark.

[0202] The amount of immobilized enzyme used is from 0.1 to 20% by weight, in particular from 10 to 15% by weight, based on the total weight of the starting materials used and to be reacted.

[0203] The inventive process is carried out at temperatures above 60° C. It is preferable to operate at temperatures of 100° C. or below that temperature. Preference is given to temperatures up to 80° C., very particular preference is given to temperatures of from 62 to 75° C., and still more preference is given to temperatures of from 65 to 75° C.

[8284] The inventive process is carried out in the pressure of a solvent. Resumples of satisfies compounds are hydrocardon of a solvent. Resumples of satisfies compounds are hydrocardon, such as possifies or a relative partial size on-leaf tenders of sevential series. Perticularly satisfies be partialled server tenders of sevential series in the form of an insome network, eighty-more, paths and the form of an insome network, eighty-more partially series of the form of an insome network, eighty-more network,

[9285] The amount of solvent added is at least 5 parts by weight, based on the weight of the starting materials used and to be reacted, preferably at least 50 parts by weight, and apprictually preferably at least 100 parts by weight, and of more than 10 000 parts by weight, of solvent are undesimble, because the reaction rate decreases materially materially lower concentrations, giving uneconomically long reaction times.

[0206] The inventive process is carried out at pressures above 500 mbar. Preference is given to the reaction at atmospheric pressure or slightly increased pressure, for example at up to 1200 mbar. It is also possible to operate under markedly increased pressure, for example at pressures up to 10 bar. The reaction at atmospheric pressure is nectioned.

[0207] The reaction time for the inventive process is usually from 4 bours to 6 days, profembly from 5 hours to 5 days, and particularly preferably from 8 hours to 4 days.

[9088] Once the reaction has ended, the high-functionality hyperbranched polyesters can be isolated, e.g. by removing the cargme by filtration and concentrating the mixture, the concentration process here usually being carried out at reduced pressure. Other work-up methods with good suitability are precipitation after addition of water, followed by washing and driving.

[6209] The high-functionality, hyperbanched polyesters obtainable by the inventive process feature particularly towards of discolored and reminfed material. For the deli-nition of hyperbanched polymers, see also F. 3. Fluxy. 1. Bit 1. Bit 1. Bit 1. Bit 1. 2000, 6, no. 1, 1-8. However, in the context of the present invention, "high-functionality hyperbanched" muons that the degree of branching, i.e. the average number of doubtried most of the context of the present invention, and the context of the present production of doubtried most of the context of the present invention, and the context of the present pr

[6210] The neventive polyesters have a molar mass M., of from 300 to 50 000 gmion, perionly by from 1000 to 20 000 gmio., particularly preferably from 1000 to 10 000 gmio. Desperionly from 1000 to 19 000 gmio. The polyelispersity is form 1.2 to 50 gmicenby from 1.40 40, particularly perferably from 1.5 to 30, and very particutarily perferably from 1.5 to 10. They are assually very scalable, i.e. clear solutions can be prepared using up to 50% by recigin, in come cases even up to 50% by weight, of the year of the control of the control of the control of the control of the particular decision of the control of the control of the control of the particles of the critical of the control of

[0211] The inventive high-functionality hyperbranched polyesters are carboxy-terminated, carboxy- and hydroxyterminated, and preferably hydroxy-terminated.

terminated, and preferably hydroxy-terminated.

[0212] The ratios of the components B1): B2) are preferably from 1:20 to 20:1, in particular from 1:15 to 15:1, and

very particularly from 1:5 to 5:1 when used in a mixture.

[0213] The inventive molding compositions may comprise, as component C), from 0 to 60% by weight, in particular up to 50% by weight, of other additives and processing aids.

[0214] The inventive molding compositions may comprise, as component C_b, from 1 to 5000 ppm, preferably from 10 to 1000 ppm, and in particular from 50 to 500 ppm, of a Ca(f) salt, preferably of a Cu(f) halide, in particular in a mixture with an alkali metal halide, preferably K1.

[0215] The monovalent copper salts used preferably comprise copper(f) acetate, copper(f) chloride, copper(f) bormide, and copper(f) indide Their amounts present correspond to from 5 to 500 ppm of copper, preferably from 10 to 250 ppm, based on polyamide.

[6216] The advantageous properties are in particular obtained if the copper has molecular dispersion within the

polyamine. This is achieved by adding, to the molding composition, a consensation comprising the polyamistic and comprising a monovalent copper and: and comprising an addin intell habile in the form of a soil, knowgeneous solution. By way of example, a typical concentrate is composed of from 19 uS95 by weight of polyamistic and from 21 to 5% by weight of a mattere composed of copper beddee or opper bounds and postsustan models. The concludes of the composed of the composed of the contraction of the

[9217] Suitable polyamides for the concentrate are homopolyamides and copolyamides, in particular nylon-6 and nylon-6,6.

[0218] The inventive molding compositions may comprise, as component O₂ from 0 to 5% by welght, perferably from 0.05% by welght, and in pattendar does 0.1 to 27% of the president data of 1 to 27% of 1 to 27%

[0219] The carboxylic acids may be monobasic or dibasic. Bxamples which may be mentioned are pelargonic acid, palmitic acid, lauric acid, marganic acid, docleanedioic acid, behenic acid, and particularly preferably stearic acid, capric acid, and also montanic acid (a mixture of fatty acids having from 30 to 40 carbon atoms).

[6220] The aliphatic alcohols may be mono- to tetrahydric. Examples of alcohols are n-butnol, n-ocianol, starryl alcohol, ethylene glycol, propylene glycol, neopentyl glycol, pentserythritol, preference being given to glycerol and pentaerythritol.

[0221] The aliphatic amines may be mono, di- or triamines. Examples of these are steera/maine, etylenendiamine, propylenediamine, beamentylenediamine, propylenediamine, particular preference being given to etylenediamine and necamentylenediamine. Correspondingly, preferred exters or amides an glyceryl disterante, may be a supplementation of the contraction of the contract

[0222] It is also possible to use mixtures of various esters or amides, or esters with amides combined, the mixing ratio here being as desired.

[0223] Examples of amounts of other usual additives C) are up to 40% by weight, preferably up to 30% by weight, of elastomeric polymers (also often termed impact modifiers, elastomers, or rubbors).

[0224] These are very generally copolymers which have preferably been built up from at least two of the following monomers: ethylene, propylene, butalene, isobutene, isoprene, chloroprene, vinyl acetate, styrene, scrylonitrile and scrylates and/or methaceylates having from 1 to 18 carbon atoms in the alcohol component.

[0225] Polymers of this type are described, for example, in Houben-Weyl, Methoden der organischen Chemie, Vol. 14/1 (Georg-Thieme-Verlag, Stuttgart, Germany, 1961), pages 392-406, and in the monograph by C. B. Bucknall, "Toughened Plactice" (Applied Science Publishers, London, UK, 1977).

[0226] Some preferred types of such elastomers are described below.

[0227] Preferred types of such elastomers are those known as ethylene-propylene (EPPM) and ethylene-propylene-diene (EPDM) rubbers.

[0228] EPM rubbers generally have practically no recidual double bonds, whereas EPDM rubbers may have from 1 to 20 double bonds per 100 carbon atoms.

[0229] Examples which may be mentlored of diene mommer for EPDM ribbers are conjugated dienes, such as inogenee and bundlene, non-conjugated dienes have be inogenee and bundlene, non-conjugated dienes having from 5 to 25 earbon stomes, such all sep-frendelines, sylvhomes, 1,5-hexadiene, 2,5-dimethyl-1,5-hexadiene and 14-dectadene, cyclic dienes, such as cyclopendines, cyclobarson, vibrations, and such as a such as a cyclopendines, cyclopendines, such as 3-schylidene-2-conformene, 2-medially-3-endromene and 2-inopropsyl-5-sorbonness, and tricyclodienes, such as 3-methylrichycid-2-1, 210⁴7-3,4-decediene, and mixtures of these. Perfectuce is given to 1,5-lexadiene, 5-ethylideneousbenness and disject-positiation. The disser conlent of the EPDM ribbers is prefrarably from 0.5 to 50% by weight, in of the robbers.

[0230] EPM and EPDM rubbers may preferably also have been grafted with reactive carboxylic acids or with derivatives of these. Examples of these are acrylic acid, meth acrylic acid and derivatives thereof, e.g. glycidyl(meth)acrylate, and also maleic anhydride.

[6231] Copolymers of ethylene with earylic acid and/or mediacrylic acid, and/or with the estern of these ecids are another group of preferred rubbers. The rubbers may also comprise describoylic acids, such as maleic acid and fumaric acid, or derivatives of these acids, e.g. esters and analystifies, and/or monomers comprising eroby groups. These monomers comprising identivelyic acid derivatives or coupsiting eyon groups are preferred by incorporated and comprising office and the compression of the compression of comprising distribuyile acid groups as the comprising of the compression of the compression of the compression of the comprising distribuyile acid groups as the compression of the comprising distribuyile acid groups as the compression of the compression of

$$E_1$$
C(COOS₂)=C(COOS₂) E_4 (I)

where R^1 to R^9 are hydrogen or alkyl groups having from 1 to 6 earbon atoms, and m is a whole number from 0 to 20, g is a whole number from 0 to 10 and p is a whole number from 0 to 5.

[0232] R¹ to R⁹ are preferably hydrogen, where m is 0 or 1 and g is 1. The corresponding compounds are malcic acid, immaric acid, malcic anhydride, allyl glycidyl ether and vinyl glycidyl ether.

[9233] Proferred compounds of the formulae I, II and I/or matter each, makes making the finate may be and untuit pacyfuls at (I/or matter each, makes makes). The comparising grown genus, such as glycidyl acrylate and glycidyl methacylate, and the esters with setterity alcohols, such tert-hutyl acrylate. Although the latter have no free carbony groups, their behavior approximates to that of the first charge and they are therefore termed monomers with latent carbony STROMS.

[0234] The copolymers are advantageously composed of from 50 to 98% by weight of etilydece, from 0.1 to 20% by weight of monomers comprising epoxy groups and/or methacrylic acid and/or monomers comprising anhydride groups, the remaining amount being (meth)acrylitets.

[9235] Particular preference is given to copolymers composed of

from 50 to 98% by weight, in particular from 55 to 95% by weight, of othylene,

from 0.1 to 40% by weight, in particular from 0.3 to 20% by weight, of glycidyl acrylate and/or glycidyl methacrylate, (meth)acrylic acid and/or maleic anhydride, and

from 1 to 45% by weight, in particular from 10 to 40% by weight, of n-butyl acrylate and/or 2-ethylhexyl acrylate.

[0236] Other preferred (meth)acrylates are the methyl.

ethyl, propyl, isobutyl and tert-butyl esters.

[0237] Besides these, comonomers which may be used are

vinyl esters and vinyl ethers.

[8238] The ethylene copolymers described above may be prepared by processes known per so, preferably by random copolymerization at high pressure and elevated temperature.

Appropriate processes are well-known.

[6239] Other preferred elastomers are emulsion polymers whose preparation is described, for example, by Blackley in the monograph "Emulsion Polymerization". The emulsiliers and catalysts which can be used are known per se.

[0240] In principle it is possible to use homogeneously structured elastomers or else those with a shell structure. The shell-type structure is determined by the sequence of addition of the individual monomers. The morphology of the polymers is advolved an end of addition.

19241 Monomers which may be mentioned from, menty as examples, for the preparation of the rubbor fraction of the elastoness are explaites, such as n-butyl scrylate and 2-sheavy arraylate, corresponding methacrytates, butdene and siopenes, and also mixtures of these. These monomers may be copolymerized with other romoners, such as stynesty to the contract of the co

[0242] The soft or rubber phase (with a glass transition temperature of below 0° C.) of the clastomers may be the core, the outer envelope or an intermediate shell (in the case of clastomers whose structure has more than two shells). Elastomers having more than one shell may also have more than one shell composed of a rubber phase.

[8243]. If one or more hard components (with glass transition temperatures above 20°C) are involved, besides the rubber planse, in the structure of the elastomer, these are generally perspected by polymerizing, as principal monomenes, styreon, suryicontrile, methacrylondirile, carebhylstyreon, penethylstyreon, or activates or methacrylines, such as methacrylondirile, and as the component of the

[0244] It has proven advantageous in some cases to use emulsion polymers which have reactive groups at their surfaces. Examples of groups of this type are epoxy, carboxy, latent carboxy, amino and amide groups, and also functional groups which may be introduced by concomitant use of monomers of the general formula

where:

[0245] R^{10} is hydrogen or C_1 - C_4 -alkyl,

[0246] R¹¹ is hydrogen or C₁·C₃-alkyl or aryl, in particular phenyl.

[0247]
$$\mathbb{R}^{12}$$
 is hydrogen, $C_1\text{-}C_{10}\text{-alkyl}$, $C_6\text{-}C_{12}\text{-aryl}$ or --OR^{15}

[6248] R¹³ is C₁-C₆-alkyl or C₆-C₁₂-aryl, if desired with substitution by O- or N-comprising groups,

[0249] X is a chemical bond or C_1 - C_{10} -alkylene or C_0 - C_{12} -arylene, or

[0250] Y is O-Z or NH-Z, and

[0251] Z is C1-C10 alkylene or C0-C12 arylene.

[0252] The graft monomers described in EP-A 208 187 are also suitable for introducing reactive groups at the surface.

[0253] Other examples which may be mentioned are actytamide, methacrylamide and substituted acrylates or methacrylates, such as (N-tert-butylamino)ethyl methacrylate, (N,N-dimethylamino)ethyl acrylate, (N,N-dimethylamino)emyl acrylate and (N,N-diethylamino)ethyl acrylate and (N,N-diethylamino)ethyl acrylate

[0254] The particles of the rubber phase may also have been crosslinked. Examples of crosslinking monomers are 1,3-butadiene, divinylbenzene, dialityl phthalate and dihydrodicyclopentadienyl acrylate, and also the compounds described in EP-A 50 265. [2825] It is also possible to the the transmerse known as grant-finding monomers, in amounters twing two or more polymerizable double bonds which react at different rates during the polymerization. Predicence is given to the use of compounds of this type in which at least our cractive group which the prediction of th

[02.56] Examples of graft-linking monomers of this type are monomers comprising ally groups, in particularly selects of onlyelically unsaturated carboxylic acids, for example ally scryotas, ally metastrapida, diality imalent, diality limmarete and diality linconate, and the corresponding monosityl components of these detachoxylic acids. Besides monomers. For further details reference may be made here, for example, by U.S. Pat. No. 4,148,366.

[0257] The proportion of these crosslinking monomers in the impact-modifying polymer is generally up to 5% by weight, preferably not more than 3% by weight, based on the impact-modifying polymers.

[0258] Some preferred emulsion polymers are listed below. Mention may first be made here of graft polymers with a core and with at least one outer shell, and having the following structure:

Type	Monomers for the core	Meacmers for the eavelope				
1	1,3-butsdiene, iroprene, a-butyl acrylate, ethythexyl acrylate, or a mixture of these	styrene, scrylonitrile, methyl methocrylete				
п	as I, but with concomitant use of crosslinking agents	as I				
ш	as I or II	n-batyl scryiste, ethyl scryiste, methyl scryiste, 1,3 butediese, isogram, ethylhexyl scryiste				
IV	as I or II	as I or III, but with concomitant use of monomers having reactive eroses, as described herein				
v	styrene, acrylonitrile, methyl methacrylate, or a mixture of these	first anvelope composed of monomers as described under I and II for the core, second envelope as described under I or IV for the envelope				

[9259] Instead of graft polymers whose structure has more than one shall, it is also possible to use honogeneous, is single-shell, classioners composed of 1,3-batdeine, isoprune and n-butyl acrylate or from copolymers of these. These products, too, may be prepared by concomitant use of crosslinking monomers or of monomers inving reactive rooms.

[0260] Examples of preferred emilsion polymers are n-butyl acrylate-(meth)scrylic acid copolymers, n-butyl acrylate-(prefly scrylate or n-butyl acrylate-(prefly scrylate or n-butyl acrylate or p-butyl acrylate or composed of n-butyl acrylate or based on butsdiene and with an outer envelope composed of the above-mentioned

copolymers, and copolymers of ethylene with comonomers which supply reactive groups

[0261] The elastomers described may also be prepared by other conventional processes, e.g. by suspension polymerization.

[0262] Preference is also given to silicone rubbers, as described in DB-A 37 25 576, BP-A 235 690, DB-A 38 00 603 and EP-A 319 290.

[0263] It is, of course, also possible to use mixtures of the types of rubber listed above.

[0264] Fibrous or particulate fillers C) which may be mentioned are carbon fibers, glass fibers, glass beads, amorphous silies, celcium melitace, enderium metalicate, magnesium carbonate, kaolin, chalk, powdered quartz, mics, barium sulfate and feldapar, used in amounts of up to 50% by weight, in particular up to 40% by weight, in particular up to 40% by weight, in

[0265] Preferred fibrous fillers which may be mentioned are carbon fibers, armid fibers and potassium timate fibers, and particular preference is given to glass fibers in the florm of E glass. These may be used as rovings or in the commercially available forms of chopped glass.

[0266] The fibrous fillers may have been surface-pretreated with a silane compound to improve compatibility with the thermoplastic.

[0267] Suitable silane compounds have the general formula:

$$(X-(CH_2)_n)_k$$
—Si-(O-C_{ss}H_{2m+1})_{4-k} where:

[0268] X is

 $\begin{bmatrix} 0269 \end{bmatrix}$ n is a whole number from 2 to 10, preferably 3 to

[0270] m is a whole number from 1 to 5, preferably 1 to

[0271] k is a whole number from 1 to 3, preferably 1.

[0272] Preferred silaae compounds are aminopropyltrimethoxysilane, aminobulyltrimethoxysilane, aminopropyltriethoxysilane and aminobulyltriethoxysilane, and also the corresponding silanes which comprise a glycidyl group as substituer X

[0273] The amounts of the silane compounds generally used for surface-coating are from 0.05 to 5% by weight, perferably from 0.5 to 1.5% by weight and in particular from 0.8 to 1% by weight (based on C).

[6274] Acicular mineral fillers are also suitable.

[0275] For the purposes of the invention, acicular mineral fillers are mineral fillers with strongly developed acicular character. An example is selectar wollastonite. The mineral preferably has an UD (depith to diameter) ratio of from 8:1 to 35:1, preferably from 8:1 to 11:1. The mineral filler may if appropriate, have been pretreated with the abuvenment tioned silner compounds, but the pretreatment is not essen[6276] Other fillens which may be mentioned are leadin, cachined localin, wouldnessine, it and chell, and also lamelar or accident nearing, wouldnessine, it as an other length of the purpose of the leading of the leading of the leading lamb of the lamb of the lamb of la

[9277] As component C), the thermseplastic molding compositions of the invention may comprise usual processing sids, such as stabilizers, oxidation retarders, agents to comtemat decomposition due to be lead and decomposition due to ultraviolet light, lubricants and mold-release agents, colorants, such as dyes and pigments, mucleating agents, plasticizers. filance retardants, etc.

[0278] Examples which may be mentioned of oxidation retarders and heat stabilizers are sterically hindered phenols and/or phosphites and amines (e.g. TAD), hydroglamms, aromatic secondary amines, such as diphenylamines, various substituted members of these groups, and mixtures of these in concentrations of up to 1% by weight, based on the weight of the thermoplastic modifies composition.

[0279] UV stabilizers which may be mentioned, and are generally used in amounts of up to 2% by weight, based on the molding composition, are various substituted resorcinols, salicylates, benzotriazoles, and benzophenones.

[0280] Colorants which may be added are inorganic pigments, such as titanium dioxide, ultramarine blue, iron oxide, and carbon black; and also organic pigments, such as pithelicoyanines, quinacridones and perylenes, and also dues such as nigrosine and anthracultones.

[6281] Nucleating agents which may be used are sodium phenylphosphinate, alumina, silica, and preferably talc.

[9282] Other Intricants and mold-releuse agents are unally used in amounts of up to 1% by weight. Perference is given to long-chain furly acids (e.g., stentic acid or behavior, acids, last of these (e.g., acidum is consist or rime stenents) or montan waxes (mixtures of straight-chain saturated carboylic acids height acid in length of from 28 to 3% carbon low-molecular veight polyethylene waxes or low-molecular-weight polyethylene waxes or low-molecular-weight polyethylene waxes.

[9283] The inventive thermoplastic molding compositions may be prepared by methods known per se, by mixing the starting compositions as sower extracted, Bibmoder mixing or paperatus, such as sower extracted, Bibmoder mixing or Bibmody mixes, and then extracting them. The extracted may then be cooled components and then to add the renatining starting materials individually and/or likewise in a mixture. The mixing temperatures are generally from 230 to 320° C.

[10244] In austher preferred procedure, components B; and, if appropriate, C may be mixed with a propolymer, and, if appropriate, C may be mixed with a propolymer, and include the consideration of the control of the

[6285] In particular, the individual components can be proceesed without difficulty (without clumping or caking)

and in short cycle times, so that a particular possible application thereof is thin-wall components.

[0286] Improved-flow polyamides could be used in almost any injection-molding application. Improved flow allows lower melt temperature and can therefore lead to a marked reduction in the overall cycle time for injection molding (reduction in production costs for an injection molding!) Furthermore, the injection pressures needed during processing are lower, thus requiring lower total locking force on the injection mold (less capital expenditure on the injectionmolding machine).

[0287] Alongside the improvements in the injection-mold-ing process, lowering of melt viscosity can lead to significant advantages in the actual design of the component. For example, injection molding can be used to produce thinwalled applications which, by way of example, were not hitherto capable of production using filled grades of polyamide. Similarly, by using grades of polyamide which are reinforced but relatively free-flowing could reduce wall thicknesses in existing applications, and therefore reduce component weight.

[0288] These materials are suitable for production of fibers, foils, or moldings of any type. Some examples are given below:

[0289] Improved-flow polyamides can be used in the electrical and electronics sector to produce plugs, plug components, plug connectors, cable harness components, circuit mounts, circuit-mount components, three-dimension-ally injection-molded circuit mounts, electrical connector elements, or mechatronic components.

[0290] Possible uses in automobile interiors are dashboards, steering-column switches, seat components, headrests, center consoles, gearbox components, and door mod-ules, and possible uses in automobile exteriors are door handles, exterior-mirror components, windshield-washer components, windshield-washer protective housings, grills, roof reils, sunroof frames, engine covers, cylinder-head covers, inlet manifolds, windshield wipers, and exterior bodywork parts.

[0291] In the kitchen and household sector, improved-flow polyamides can be used to produce components for kitchen machines, e.g. fryers, smoothing irons, buttons, and applications in the garden and leisure sector, e.g. components for irrigation systems or garden machines, and door handles.

EXAMPLES

[0292] The following components were used:

Component A/1:

[0293] Nylon-6 (polycaprolactam) with a viscosity number VN of 150 ml/g, measured in the form of a 0.5% strength by weight solution in 96% strength by weight sulfuric acid at 25° C. to ISO 307 (the material used being Ultramid® B3 from BASF AG).

Component A/2

[0294] Nylon-6T/6 (70:30) with a viscosity number VN of 135 ml/g, measured in the form of a 0.5% strength by weight solution in 96% strength by weight sulfuric acid at 25° C. to ISO 307 (the material used being Ultramid® KTR 4350 from BASF AG). Component B

[0295] Polycarbonates B1

[0296] General operating specification:

[0297] The polyhydric alcohol, diethyl carbonate, and catalyst (250 ppm, based on weight of alcohol) were used as initial charge in a three-necked flask equipped with stirrer, reflux condenser, and internal thermometer, the amounts in the mixture being as in table 1, and the mixture was heated to 120° C. and stirred at this temperature for 2 h. The temperature of the reaction mixture reduced here as the reaction proceeded, because of onset of evaporative cooling by the ethanol liberated. The reflux condenser was then replaced by an inclined condenser and, based on the equivalent amount of estalyst, one equivalent of phosphoric acid was added, the ethanol was removed by distillation, and the temperature of the reaction mixture was increased slowly to 160° C.

[9298] The alcohol removed by distillation was collected in a cooled round-bottomed flask and weighed, and conversion was thus determined and compared in percentage terms with the full conversion theoretically possible (see table 1).

[0299] The reaction products were then analyzed by gel permeation chromatography, using dimethylacetamide as eluent and polymethyl methacrylate (PMMA) as standard.

[9366] Glass transition temperature and, respectively, melting point were determined by DSC (differential scanning calorimetry) to ASTM 3418/82, evaluating the second heating curve.

TABLE 1

Comp.	Alcohol	Alcohol:Diethyl carbonate molar ratio	Catalyst	Distillate, amount of alcohol, based on full convenion Mol %	Molar mess of product (gined) Mw Mn	Vise, of product at 23° C. (mPss)	OH number of product to THN 532MO, Part 2 (mg KOH/g)
B 1/1	TMP × 3 EO	1:1	K2CO3	90	4600	5030	310
B 1/2	IMP × 1.2 PO	1:1	K_2CO_3	90	2600 2800 1800	37 000	436
B 1/3	TMP × 12 EO	1.1	KOH	70	4400	550	189
В 3/4	TMP \times 5.4 PO	1:1	кон	85	2500 4000 1700	1900	228

TMP = Trimethylolprope

EO - Ethylene oxide

PO - Propylene oxide

Component B 2/1

[6891] 2001 g (1.5 mol) of cyclohexame-1,2-disarbraylte antilyrdine and 979 (1.6 mol) of glycerol were used a mittale charge in a 4 i jacketed reactor equipped with stirrer, internal thermomenter, go-inder thee, Polito, coucleurer, and vacuum connection with cold trap. After addition of 3 g of disab-strylin coids, the mixture was heated to an internal temperature of 150° C, with the aid of an oil beth, Archaecel pressure of 140 mbr was applied in order to remove water formed during the reaction. The reaction mixture was kept at the stated temperature and the stated pressure for 120 mon. Cooling to room temperature gave the solid polyseiter. The analytical data are given in table 2.

Component B 2/2

[6042] On [6041 mot) of edipic noid and 89 g (0.34 mot) of trichydrocynophilocynamics (HIRC) were used as initial charge in a 25 ml flow-necked glass flask equipped with sizre, internal flemomente, gar-hiet thee, return consideration and vacuum connection with ords large. After motion of the control of th

Component B 2/3

[BMS] II h.2 § (M.o. ma)) of dimethyl suspitationist (JMT) and 335 § (0.5 no) of ethoxylated trilighydxoxymothylphopane, moline mass 670 g/mol, were used as initial charge in a 11 reaction, vested equipped with streng; internal thermomenter, gas-linkt tube, reflex rostediences, and vested month consecution of the strength of the str

TABLE 2

No.	Mn (g/ mol)	Mw (g/mol)	Tg (* C.)	Acid number (mg KOH/g)	Hydroxy mamber (mg KOH/g)	Viscosity (mPa·s)
B 2/1	1280	3310	26.5	141	268	700 @ 150° C.
B 2/2	4160	23 770	n.d.	n.d.	n.d.	5000 @ 50° C.
15 2/3	11.30	3030	15.4	37	369	5400 48 100° C.

EGS - IKS CARGINING

Analysis of Inventive Polyesters

[0304] The polyesters were analyzed by gel permeation chromatography, using a refractometer as detector. The mobile phase used was tetrahydrofuran and the standard used to determine molecular weight was polymethyl methacytate (PMMA). [0305] Glass transition temperatures were determined by differential scanning calorimetry (D8C) evaluating the scoond heating curve. Acid number and hydroxy number were determined to DIN 53240, Part 2.

Preparation of Molding Compositions for Constitutions with A/3

[0306] Components A) and B) were blended at from 250 to 260° C. in a twin-screw extruder and extruded into a water bettermined on the pelletization and drying, VN and MVR were determined on the pellets.

Preparation of Molding Compositions for A/2

[0307] Components A) and B) were blended at from 320 to 330° C. in a twin-screw extruder and extruded into a water bath. After pelletization and drying, VN and MVR were determined on the pellets

[0308] MVR was determined to ISO 11 33 (at 275° C., melting time: 8 min, 5 kg), and for A/2 at 325° C., melting time: 4 min., and 5 kg, and VN was determined to ISO 307 (at c-5 c) in 95% strength culture ooid).

[0309] The variables E, σ , ϵ , ak, and an were measured to ISO 527 and 179 1 eA, and also 179 1 eV.

[6310] The inventive constitutions and the results of the measurements are found in tables 3 to 5.

TARLE 3

	Component A/1 [% by weight]	Component B [% by weight]	VN [ml/g]	MVR [ml/10 min]	
Comparison	100	0	150	131	
Ex. I	96	4% B I/I	144	236	
Ex. 2	99	1% B 1/2	148	203	
Ex. 3	98	2% B I/3	149	193	
Ex. 4	99	1% B 2/1	144	272	
Ex. 5	99	1% B 2/2	144	197	
Ex. 6	99	1% B 2/3	147	208	

[0311]

TABLE 4

	Com- powent A/2 [% by weight]	Com- ponent B [% by weight]	VN [ml/g]	MVR [ml/10 min]	
Comparison	100	0	126	52.3	
Ex. 1	99	1% B 1/2	123	67.4	
Es. 2	98	2% D 1/2	123	62.6	
Ex. 3	99	1% B 1/4	126	55.4	
Ex. 4	98	2% B 1/4	122	77.2	

「**0312**1

TARLE S

	Component A/2 [% by wt.]	Component B	Flow spiral 80/330 cm	Flow spiral 100/330 om	B MPn	o MPa	e %	ak kVm²	sa: kU/m²
Comparison	100	0	42	46	3170	95	3.9/	8,8	171
Ex. 1	99	154 B 54	46	49	3160	94	3.6/5.1	8.8	276
Ex. 2	96	2% B 15	50	52.5	3200	95	4.2/5.1	8.4	211
Ex. 3	99	1% B ¼	45.5	49	3120	95	4.4/5.2	9.3	137
Ex. 4	98	256 B 54	48.5	59	3170	90	3.7/3.7	8,8	46

80/330 Mold temperature/readuct temperature

1. A thermoplastic molding composition, comprising:

 A) from 10 to 99% by weight of at least one thermoplastic polyamide;

B) from 0.01 to 50% by weight of

Bi) at least one highly branched or hyperbranched polycarbonate with an OH number of from 1 to 600 mg KOH/g of polycarbonate (DIN 53240, Part 2), or

B2) at least one highly branched or hyperbranched polyester of $A_x B_y$ type, where x is at least 1.1 and y is at least 2.1, or a mixture of these; and

C) from 0 to 60% by weight of other additives;
 wherein the total of the percentages by weight of com-

ponents A) to C) is 100%.

2. The thermoplastic molding composition according to

claim 1, wherein component B1) has a number-average molar mass Mn of from 100 to 15 000 g/mol. 3. The thermoplastic molding composition according to claim 1, wherein component B1) has a glass transition

temperature Tg of from -80° C. to 140° C.

4. The thermoplastic molding composition according to claim I, wherein component B1) has a viscosity (mPas) at

claim 1, wherein component B1) has a viscosity (mPas) at 23° C. (DIN 53019) of from 50 to 200 000.

5. The thermoplastic molding composition according to claim 1, wherein component D2) has a number-average molar mass Mn of from 300 to 30 000 g/mol.
 6. The thermoplastic molding composition according to

claim 1, component B2) has a glass transition temperature Tg of from -50 to 140° C.

7. The thermoplastic molding composition according to

claim 1, wherein component B2) has an OH number (DIN 53240) of from 0 to 600 mg KOH/g of polyester.

8. The thermoplastic molding composition according to

claim 1, wherein component B2) has a COOH number (DIN 53240) of from 0 to 600 mg KOH/g of polyester.

9. The thermoplastic molding composition according to

 The thermoplastic molding composition according to claim 1, wherein component B2) has at least one OH number or COOH number greater than 0. The thermoplastic molding composition according to claim 1, wherein the ratio of components B1):B2) is from 1:20 to 20:1.

11. (canceled)

 A fiber, a foil, or a molding, obtainable from the thermoplastic molding composition according to claim 1.
 A method of making a fiber, foil, or molding, the

method comprising:

preparing a thermoplastic molding composition according
to claim 1; and

to claim 1; and forming a fiber, foil, or molding from the thermonlastic

forming a fiber, foil, or molding from the thermoplastic molding composition.

 The thermoplastic molding composition according to claim 2, wherein component B1) has a glass transition temperature Tg of from -80° C. to 140° C.

15. The thermoplastic molding composition according to claim 2, wherein component B1) has a viscosity (mPas) at 23° C. of from 50 to 200 000.

16. The thermoplastic molding composition according to claim 3, wherein component B1) has a viscosity (mPas) at 23° C. of from 50 to 200 000.

17. The thermoplastic molding composition according to claim 2, wherein component B2) has a number-average molar mass Mn of from 300 to 30 000 g/mol.

18. The thermoplastic molding composition according to claim 3, wherein component B2) has a number-everage

molar mass Mn of from 300 to 30 000 g/mol.

19. The thermoplastic molding composition according to claim 4, wherein component B2) has a number-average

molar mass Mn of from 300 to 30 000 g/mol.

20. The thermoplastic molding composition according to claim 2, wherein component B2) has a glass transition

temperature Tg of from -50 to 140° C.

21. The thermoplastic molding composition according to claim 3, wherein component B2) has a glass transition temperature Tg of from -50 to 140° C.

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